FOOD AND DRUG ADMINISTRATION (FDA) Center for Drug Evaluation and Research (CDER)

Arthritis Advisory Committee (AAC) Meeting

FDA White Oak Campus, Building 31 Conference Center, the Great Room (Rm. 1503) 10903 New Hampshire Avenue, Silver Spring, Maryland July 13, 2016

DRAFT AGENDA

The committee will discuss biologics license application (BLA) 761042, for GP2015, a proposed biosimilar to Amgen Inc's Enbrel (etanercept) submitted by Sandoz, Inc. The proposed indications (uses) for this product are (1) Reducing signs and symptoms, inducing major clinical response, inhibiting the progression of structural damage, and improving physical function in patients with moderately to severely active rheumatoid arthritis (RA)((in combination with methotrexate (MTX) or used alone); (2) reducing signs and symptoms of moderately to severely active polyarticular juvenile idiopathic arthritis in patients ages 2 and older; (3) reducing signs and symptoms, inhibiting the progression of structural damage of active arthritis, and improving physical function in patients with psoriatic arthritis (in combination with MTX in patients who do not respond adequately to MTX alone); (4) reducing signs and symptoms in patients with active ankylosing spondylitis; (5) treatment of adult patients (18 years or older) with chronic moderate to severe plaque psoriasis who are candidates for systemic therapy or phototherapy.

7:30 a.m.	Call to Order and Introduction of Committee	Daniel Solomon, MD, MPH Acting Chairperson, AAC
7:35 a.m.	Conflict of Interest Statement	Moon Hee Choi, PharmD Acting Designated Federal Officer, AAC
7:40 a.m.	351(k) Regulatory Pathway	Leah Christl, PhD Associate Director, Therapeutic Biologics Therapeutic Biologics and Biosimilars Staff Office of New Drugs (OND), CDER, FDA
8:20 a.m.	Clarifying Questions to the FDA	
8:25 a.m.	FDA Introductory Remarks	Nikolay P. Nikolov, MD Clinical Team Leader Division of Pulmonary, Allergy & Rheumatology Products (DPARP) Office of Drug Evaluation II (ODE-II) OND, CDER, FDA
8:30 a.m.	APPLICANT PRESENTATIONS	Sandoz, Inc.
	Introduction and Concept	Mark McCamish, MD, PhD Global Head of Development Sandoz Biopharmaceuticals
	Analytical Demonstration of Similarity	Martin Schiestl, PhD Chief Science Officer, Sandoz Biopharmaceuticals
	Non-clinical and Pharmacokinetic Characterization of GP2015	Oliver von Richter, PhD, FCP Clinical Pharmacologist, Global Clinical Development Sandoz Biopharmaceuticals

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DRAFT AGENDA (cont.)

APPLICANT PRESENTATIONS (co	ont.)
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Clinical Confirmation of GP2015

Equivalence to Enbrel®

Use in Clinical Practice

Conclusions

10:15 a.m. **BREAK**

10:00 a.m.

10:30 a.m. **FDA PRESENTATIONS**

GP2015 Product Quality Review

Clarifying Questions to Applicant

GP2015 Statistical Equivalence

Testing for Bioactivity

Clinical Pharmacology Review

Clinical Efficacy Review

Clinical Safety and Immunogenicity Review, Considerations for Extrapolation and Summary of FDA Presentation Malte Peters, MD

Global Head of Clinical Development

Sandoz Biopharmaceuticals

Jonathan Kay, MD

Timothy S. and Elaine L. Peterson Chair in Rheumatology

Professor of Medicine

Director of Clinical Research, Rheumatology University of Massachusetts Medical School

Mark McCamish, MD, PhD

Peter L. Adams, PhD

CMC Product Quality Reviewer

Division of Biotechnology Review and Research 1

Office of Biotechnology Products

Office of Pharmaceutical Quality, CDER, FDA

Meiyu Shen, PhD

CMC Statistical Reviewer Division of Biometrics VI Office of Biostatistics (OB)

Office of Translational Sciences (OTS), CDER, FDA

Yunzhao Ren, PhD

Clinical Pharmacology Reviewer Division of Clinical Pharmacology II

Office of Clinical Pharmacology, OTS, CDER, FDA

Kathleen Fritsch, PhD

Mathematical Statistician

Division of Biometrics III, OB, OTS, CDER, FDA

Rachel Glaser, MD

Medical Officer

DPARP, ODE-II, OND, CDER, FDA

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DRAFT AGENDA (cont.)

12:00 p.m.	Clarifying Questions to FDA	
12:15 a.m.	LUNCH	
1:15 p.m.	OPEN PUBLIC HEARING	
2:45 p.m.	Charge to the Committee	Nikolay P. Nikolov, MD
3:00 p.m.	Questions to the Committee and Committee Discussion	
3:30 p.m.	Break	,
3:45 p.m.	Questions to the Committee and Committee Discussion	
5:00 p.m.	ADJOURNMENT	